AMENDMENT TO THE CLAIMS

This listing of claims will replace all prior versions, and listings of claims in the application.

1. (Currently amended) A compound of formula I.

wherein:

R1 and R4 are each, independently,

H;

 $C_{1^{*}}C_{11^{*}}$ alkyl, $C_{2^{*}}C_{10^{*}}$ alkenyl or $C_{2^{*}}C_{10^{*}}$ alkynyl, each of which is optionally substituted one or more times by F, OH. $C_{1^{*}}C_{3^{*}}$ alkoxy, $C_{1^{*}}C_{3^{*}}$ alkylmercapto, -CN, COOR⁶, CONR⁷R⁸, phenyl or heteroaryl, wherein the phenyl and heteroaryl are each independently optionally substituted one or more times by halogen, -CN, $C_{1^{*}}C_{3^{*}}$ alkyl, $C_{1^{*}}C_{3^{*}}$ alkoxy or $C_{1^{*}}$ 3.

phenyl or heteroaryl, each of which is optionally substituted one or more times by halogen, -CN, C_3 -C₃-alkyl, C_4 -C₄-alkoxy or CF₅:

COR 6:

CONR 10R11:

COOR 12;

CFu

halogen;

-CN:

NR 13R 14:

OR 15.

S(O)...R16:

SO₂NR¹⁷R¹⁸; or

NO2;

R2 and R3 are each, independently,

11.

halogen;

-CN:

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C:-Cir-alkyl, optionally substituted one or more times by OH, phenyl, or heterogryl:
        OH:
        C1-C10-alkoxy;
        phenoxy;
        S(O),,R19;
        CF<sub>3</sub>.
        NOs:
        C1-C10-alkylamino:
        di(C<sub>1</sub>-C<sub>10</sub>-alkyl)amino;
        (C<sub>1</sub>-C<sub>6</sub>-alkyl)-CONH-:
        phenyl-CONH- or phenyl-SO<sub>2</sub>-O-, wherein the phenyl is optionally substituted one or more times by
        halogen, -CN, methyl or methoxy;
        Ci-Ca-alkyl-SO5-O-:
        (C1-C6-alkyl)-CO-, wherein the C1-C6-alkyl is optionally substituted one or more times by F, di(C1-C3-
        alkyl)amino, pyrrolidinyl or piperidinyl; or
        phenyl-CO-, wherein the phenyl is optionally substituted one or more times by Ci-Ci-alkyl, halogen
        or methoxy:
Rs is Ar or Hetar, each of indolyl which is optionally substituted one or more times by
                 halogen;
                 -CN;
                 NH<sub>2</sub>;
                 C1-C10-alkvi, C2-C10-alkenvi, C2-C10-alkvnvi, C1-C10-alkoxv, C1-C10-alkviamino or di(C1-C10-
                 alkyl)amino, wherein the alkyl, alkenyl, alkynyl and alkoxy are each independently optionally
                 substituted one or more times by F, OH, C1-C8-alkoxy, aryloxy, C1-C8-alkylmercapto, NHs.
                 C1-C8-alkylamino or di(C1-C8-alkyl)amino;
                 C3-C5-alkandiyl;
                 phenyl:
                heteroaryl;
                 arvi-substituted or heteroarvi-substituted C.-C.-alkvi:
                CFs:
                NOs:
                 OH:
                phenoxy;
                benzyloxy;
                (C<sub>1</sub>-C<sub>i0</sub>-alkyl)-COO-;
                S(O)...R20:
                SH:
                phenylamino:
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benzylamino:
(C1-C10-alkyl)-CONH-;
(C1-C10-alkyl)-CO-N(C1-C4-alkyl)-;
phenyl-CONH-;
phenyl-CO-N(C1-C4-alkyl)-:
heteroaryl-CONH-;
heteroaryl-CO-N(C1-C4-alkyl)-:
(C<sub>1</sub>-C<sub>10</sub>-alkyl)-CO-;
phenyl-CO-;
heteroaryl-CO-;
CF<sub>3</sub>-CO-:
-OCH-O-:
-OCF-O-:
-OCH2CH2O-:
-CH<sub>2</sub>CH<sub>2</sub>O-:
COOR21:
CONR 22 R 23.
CONH)-NH-:
SO<sub>2</sub>NR<sup>24</sup>R<sup>25</sup>:
R36SO-NH-:
R27SO-N(C1-C6-alkyl)-; or
a residue of a saturated or unsaturated aliphatic, monocyclic 5-membered to 7-membered
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heterocycle containing 1, 2 or 3 heteroatoms selected from the group consisting of N, O and S, wherein the heterocycle is optionally substituted one or more times by halogen, C₁-C₃-alkyl, C₁-C₃-alkoxy, OH, oxo or CF₅, and the heterocycle is optionally condensed to the group Ar or the group Heterindolyl group:

wherein all aryl, heteroaryl, phenyl, aryl-containing, heteroaryl-containing and phenyl-containing groups, which are optionally present in the said substituents of the said-group Ar or the said group Hetarindolyl group, can be substituted by one or more substituents selected from the group consisting of halogens, -CN. C₁-C₂-alkyl, OH, C₁-C₂-alkoxy, and CF₃;

Rois F

 $C_{t^*}C_{t^*}$ -alkyl, optionally substituted one or more times by F, $C_{t^*}C_{t^*}$ -alkoxy or di($C_{t^*}C_{t^*}$ -alkyl) arnino; aryl- $(C_{t^*}C_{t^*}$ -alkyl)- or heteroaryl- $(C_{t^*}C_{t^*}$ -alkyl)- either of which is optionally substituted one or more times by halogen, $C_{t^*}C_{t^*}$ -alkyl, $C_{t^*}C_{t^*}$ -alkoxy or di($C_{t^*}C_{t^*}$ -alkyl)-amino;

R⁷ is H;

 $C_{1^{*}}C_{10^{*}}$ alkyl, optionally substituted one or more times by $F, C_{1^{*}}C_{3^{*}}$ alkoxy, $di(C_{1^{*}}C_{3^{*}}$ alkyl) antino or phenyl; or

pbenyl, indanyl or heteroaryl, each of which is optionally substituted one or more times by halogen, -CN, C_1-C_2 -alkyl, C_1-C_3 -alkyxy or CF_3 ;

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R8 is H or C1-C10-alkyl;
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R⁹ is C₁-C₁-galkyl, optionally substituted one or more times by F, C₂-c₄-alkoxy or di(C₂-C₂-alkyl)amino; or phenyl or heteroaryl, each of which is optionally substituted one or more times by C₁-C₂-alkyl, C₁-C₂alkoxy, balogen, -CN or CF₅;

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R<sup>10</sup>, independently from R<sup>7</sup>, is R<sup>7</sup>:
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R1, independently from R8, is R8;

R12, independently from R6, is R6;

R^D is H:

C1-C6-alkyl; or

phenyl, benzyl, heteroaryl, (C₁-C₆-alkyl)-CO-, phenyl-CO-, or heteroaryl-CO-, each of which is optionally substituted one or more times by halogen, -CN, C₁-C₃-alkyl, C₁-C₃-alkoxy or CF₅;

R14, independently from R13, is R13;

R15 is H:

C1-Cm-alkyl;

(C₁-C₂-alkoxy)-C₁-C₂-alkyl-;

benzyl, phenyl or heteroaryl, each of which is optionally substituted one or more times by halogen.

-CN, C₁-C₁-alkyl, C₁-C₂-alkoxy or CF₃:

R¹⁶ is C₁-C₁₅-alkyl, optionally substituted one or more times by F, OH, C₁-C₈-alkoxy, aryloxy, C₁-C₈-alkylmercapio, C₁-C₈-alkylamino or di(C₁-C₈-alkyl)amino;

CFe or

phenyl or heteroaryl, each of which is optionally substituted one or more times by halogen, \cdot CN, C_3 - \cdot Cl₃-alkyl, C_4 - \cdot Cl₃-alkyl, C_4 - \cdot Cl₄-alkoxy or CE₄;

R17, independently from R7, is R7;

R18, independently from R8, is R8;

R19, independently from R16, is R16.

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R20, independently from R10, is R16;
  R21, independently from R6, is R6;
  R22, independently from R7, is R7;
 R28, independently from R8, is R8;
 R31, independently from R7, is R7;
 R25, independently from R8, is R8;
 R26, independently from R16, is R16;
 R27, independently from R16, is R16;
 R<sup>30</sup> is H:
                       C1-C10-alkyl, C7-C10-alkenyl or C7-C10-alkynyl, each of which is optionally substituted one or more
                       times by F. OH, C1-C8-alkoxy, C1-C8-alkylmercapto, -CN, COOR31, CONR32R33, NR34R35, (C1-C8-alkylmercapto, -CN, COOR31, CONR32R33, (C1-C8-alkylmercapto, -CN, COOR31, COOR31, (C1-C8-alkylmercapto, -CN, COOR31, COOR31, (C1-C8-alkylmercapto, -CN, COOR31, COOR31, COOR31, (C1-C8-alkylmercapto, -CN, COOR31, COOR31, (C1-C8-alkylmercapto, -CN, COOR31, COOR31, (C1-C8-alkylmercapto, -CN, COOR31, COOR31, (C1-C8-alkylmercapto, -CN, COOR31, (C1-C8-alkylmercapto, -CN, COOR31, (C1-C8-alkylmercapto, -CN, COOR31, (C1-C8-alkylmercapto, -CN, C1-C8-alkylmercapto, -CN, C1-C8-alkylmercapto, -CN, C1-C8-alkylmercapto, -CN, C1-C8-alkylmercapto, -CN, C1-C8-alkylmercapto, -CN, C1
                       alkyl)-CONH-. (Ci-Ci-alkoxy)-CONH-, benzyloxy-CONH-, phenyl or heteroaryl, wherein the phenyl
                       and beteroaryl are each independently optionally substituted one or more times by halogen, -CN, Cr-
                       C3-alkyl, C4-C5-alkoxy or CF3; or
                       phenyl or heteroaryl, each of which is optionally substituted one or more times by halogen, -CN, C1-
                       C3-alkyl, C4-C3-alkoxy or CF3;
 R31, independently from R6, is R6;
 R32, independently from R6, is R6:
R33, independently from R6, is R6;
R34, independently from R6, is R6:
R35, independently from R6, is R6;
 X is NR30, S. O. CH=CH, N=CH or CH=N:
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heteroaryl is a residue of a 5-membered to 10-membered, aromatic, monocyclic or bicyclic heterocycle containing one or more heteroatoms selected from the group consisting of N, O and S;

the group Heter is a residue of a 5-membered to 10 membered, aromatic, monocyclic or bleyelic heterocycle containing one or more beteroatoms selected from the group consisting of N, O and S;

aryl is phenyl, naphth-1-yl or naphth-2-yl;

the group Ar is phenyl naphth 1 yl or naphth 2 yl; and

m is 0, 1 or 2:

provided that the compound is not 2-methyl-6-trifluoromethyl-1H-indole-3-carboxylic acid benzothiazol-2-ylamide;

or a stereoisomer or a mixture of stereoisomers thereof in any ratio, or a pharmaceutically acceptable salt thereof.

2-5. (Cancelled)

6. (Currently amended) AThe compound according to claim 1, wherein:

R¹ and R⁴ are each, independently,

13:

Halogen; or

C₁-C₄-alkyl;

and

R2 and R3 are each, independently,

H:

Halogen; or

Cı-Cı-alkyl.

7. (Currently amended) AThe compound according to claim 1, wherein:

Rb is phenyl or Heter, each ofindoly! which is optionally substituted one or more times by

halogen;

-CN;

NH₃:

 $C_1\text{-}C_6\text{-}alkyl, C_2\text{-}C_6\text{-}alkenyl, C_2\text{-}C_6\text{-}alkynyl, C_1\text{-}C_3\text{-}alkoxy, C_1\text{-}C_4\text{-}alkylamino or di(}C_1\text{-}C_4\text{-}alkyl)\text{amino, each of which is optionally substituted one or more times by F, }C_1\text{-}C_3\text{-}alkoxy, }$

C1-C3-alkylmercapto or NH2;

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C:-C:-alkandivl:
 phenyl;
 heteroaryl:
 phenyl-substituted or heteroaryl-substituted C1-C2-alkyl;
CF<sub>1</sub>:
OH:
(C<sub>1</sub>-C<sub>4</sub>-alkyl)-COO:
S(O)_{m^*}(C_1-C_4)-alkyl;
(C1-C4-alkyl)-CONH-;
(C1-C4-alkyl)-CON(C1-C4-alkyl)-;
(Cr-C4-alkyl)-CO-;
phenyl-CO-;
heteroaryl-CO-:
CF-CO-:
-OCH2O-;
-OCF2O-;
-OCH-CH-O-:
-CH-CH-O-:
-COO(C1-C6-alkyl);
-CONH-:
-CONH(C1-C4-alkyl);
-CON(di(C1-C1-alkyl));
-C(NH)NH2;
-50<sub>2</sub>NH<sub>2</sub>;
-SO2NH(C1-C4-alkyl);
-SO2NH(phenyl);
-SO2N(di(C1-C1-alkyl)):
(C1-C4-alkyl)-SO3NH-;
(C1-C4-alkvl)-SO-N(C1-C4-alkvl)-; or
a residue of a saturated or unsaturated aliphatic, mononuclear 5-membered to 7-membered
heterocycle containing 1, 2 or 3 heteroatoms selected from the group consisting of N, O and S.
wherein the heterocycle is optionally substituted one or more times by halogen, C1-C3-alkyl,
C1-C3-alkoxy, OH, oxo or CF3, and the heterocycle is optionally condensed to the said phenyi
or the said-group Hetarindolyl group;
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wherein all heteroaryl, phenyl, heteroaryl-containing and phenyl-containing groups, which are optionally present in the said substituents of the said phenyl or the said group Heterindolyl group, can be substituted by one or more substituents selected from the group consisting of halogen, -CN, C_1 - C_2 -alkoxy, and CF_3 -

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- (Currently amended) A pharmaceutical composition comprising a pharmaceutically effective amount of effic compound according to claim 1 and a pharmaceutically acceptable carrier.
- (Withdrawn-currently amended) A method for the stimulation of the expression of endothelial NO synthase, in a patient in need thereof, comprising administering to the patient a pharmaceutically effective amount of athe compound according to claim 1.
- 10. (Withdrawn-currently amended) A method for the treatment of cardiovascular diseases, stable or unstable angina pectoris, coronary heart disease, Prinzmetal angina, acute coronary syndrome, heart failure, myocardial infarction, stroke, thrombosis, peripheral artery occlusive disease, endothelial dysfunction, atherosclerosis, restenosis, endothel damage after PTCA, hypertension, essential hypertension, palmonary hypertension, secondary hypertension, renovascular hypertension, chronic glomerulonephritis, erectile dysfunction, ventricular arrhythmia, diabetes, diabetes complications. nephropathy, tetinopathy, angiogenesis, asthma bronchiale, chronic renal failure, cirrhosis of the liver, osteoporosis, restricted memory performance or a restricted ability to learn, or for the lowering of cardiovascular tisk of postmenopausal women or of women taking contraceptives, in a patient in need thereof, comprising administering to the patient a pharmaceutically effective amount of eithe compound according to claim 1.
- 11. (New) The compound according to claim 1, wherein

R⁵ is <u>indolyl</u> which is attached via ring carbon atom and which is optionally substituted one or more times by:

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halogen:
-CN:
NH<sub>2</sub>;
C1-C10-alkyl, C2-C10-alkenyl, C2-C10-alkynyl, C1-C10-alkoxy, C1-C10-alkylamino or di(C1-C10-
alkyl)amino, wherein the alkyl, alkenyl, alkynyl and alkoxy are each independently optionally
substituted one or more times by F, OH, C<sub>1</sub>-C<sub>8</sub>-alkoxy, aryloxy, C<sub>1</sub>-C<sub>8</sub>-alkylmercapto, NH<sub>2</sub>,
Ci-Cs-alkylamino or di(Ci-Cs-alkyl)amino;
CvCcalkandivl:
phenyl:
heteroaryl;
arvl-substituted or heteroaryl-substituted C1-C4-alkyl;
CF3;
NO<sub>2</sub>;
OH:
phenoxy;
benzyloxy;
(Ci-Cie-alkyl)-COO-:
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S(O),,R20;
SH;
phenylamino;
benzylamino:
(Ci-Cia-alkyl)-CONH-;
(C_1-C_{10}-alkyl)-CO-N(C_1-C_4-alkyl)-;
phenyl-CONH-;
phenyl-CO-N(C1-C4-alkyl)-;
heteroarvi-CONH-:
heteroaryl-CO-N(C1-C4-alkyl)-;
(C1-C10-alkyl)-CO-;
phenyl-CO-;
heteroaryl-CO-;
CF<sub>t</sub>-CO-;
-OCH-O-:
-OCF-O-:
-OCH2CH2O-;
-CH2CH2O-;
COOR21;
CONR 22 R 23.
C(NH)-NH-:
SO<sub>2</sub>NR<sup>24</sup>R<sup>25</sup>;
R36SO-NH-:
R27SO2N(C1-C6-alkyl)-; or
```

a residue of a saturated or unsaturated aliphatic, monocyclic 5-membered to 7-membered heterocycle containing 1, 2 or 3 heteroatoms selected from the group consisting of N, O and S, wherein the heterocycle is optionally substituted one or more times by halogen, Ct-C₃-

 $alkyl, C_1\hbox{-} C_3\hbox{-} alkoxy, OH, oxo or CF_3, and the heterocycle is optionally condensed to the indolyl group; }$

wherein all aryl, heteroaryl, phenyl, aryl-containing, heteroaryl-containing and phenyl-containing groups, which are optionally present in the said substituents of the indolyl group, can be substituted by one or more substituents selected from the group consisting of halogens, -CN, C₃-C₃-alkyl, OH, C₃-C₃-alkoxy, and CF₃.

12. (New) The compound according to claim 1 of formula Ik: